Uploading C:\Program Files\Stnexp\Queries\10510333.str

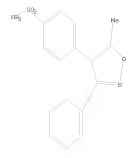
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18 19 20 ring nodes:
1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 chain bonds:
3-19 6-13 11-14 15-18 19-20 ring bonds:
1-2 1-6 2-3 3-4 4-5 5-6 7-8 7-12 8-9 9-10 10-11 11-12 13-14 13-15 14-16 respond to the second term of the
```

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:CLASS 19:CLASS 20:CLASS

L1 STRUCTURE UPLOADED

=> d L1 HAS NO ANSWERS L1 STR

chain nodes :



Structure attributes must be viewed using STN Express query preparation.

=> s 11 full

FULL SEARCH INITIATED 08:36:54 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 1043 TO ITERATE

100.0% PROCESSED 1043 ITERATIONS SEARCH TIME: 00.00.01 162 ANSWERS

L2 162 SEA SSS FUL L1

=> d 12 1-10

- L2 ANSWER 1 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN
- RN 1027903-23-6 REGISTRY
- ED Entered STN: 13 Jun 2008
- CN INDEX NAME NOT YET ASSIGNED
- MF C17 H17 N3 O6 S2
- SR Other Sources

Database: ChemSpider (ChemZoo, Inc.)

$$0 = S - NH_2$$
 OMe
$$0 = N - N$$
 OMe
$$0 = N - N$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ANSWER 2 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN

1026800-82-7 REGISTRY Entered STN: 09 Jun 2008 RN

ED

CN Benzenesulfonamide, 4-[5-methyl-3-(2,4,6-trimethoxyphenyl)-4-isoxazolyl]-(CA INDEX NAME)

C19 H20 N2 O6 S MF

Other Sources SR

Database: ChemSpider (ChemZoo, Inc.)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- L2 ANSWER 3 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN
- RN 958643-42-0 REGISTRY
- ED Entered STN: 18 Dec 2007
- CN γ-Cyclodextrin, compd. with 4-(5-methyl-3-phenyl-4isoxazolyl)benzenesulfonamide (6:1) (CA INDEX NAME)
- FS STEREOSEARCH
- C48 H80 O40 . 1/6 C16 H14 N2 O3 S MF

SR CA LC STN Files: CA, CAPLUS

CM 1

CRN 181695-72-7 CMF C16 H14 N2 O3 S

CM 2

CRN 17465-86-0 CMF C48 H80 O40

1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 4 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN RN 958643-41-9 REGISTRY

RN 958643-41-9 REGISTRY ED Entered STN: 18 Dec 2007 CN β -Cyclodextrin, compd. with 4-(5-methyl-3-phenyl-4-isoxazolyl)benzenesulfonamide (6:1) (CA INDEX NAME)

FS STEREOSEARCH

MF C42 H70 O35 . 1/6 C16 H14 N2 O3 S

SR CA

LC STN Files: CA, CAPLUS

CM 1

CRN 181695-72-7

CMF C16 H14 N2 O3 S

CM 2

CRN 7585-39-9 CMF C42 H70 O35

Absolute stereochemistry.

PAGE 1-A



1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 5 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN

RN 943600-05-3 REGISTRY

ED Entered STN: 30 Jul 2007

CN Benzenesulfonamide, 4-[3-(2-chloropheny1)-5-methyl-4-isoxazoly1]- (CA INDEX NAME)

MF C16 H13 C1 N2 O3 S

SR CA

LC STN Files: CA, CAPLUS, CASREACT

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 6 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN

RN 937250-85-6 REGISTRY

ED Entered STN: 14 Jun 2007

N Acetamide, N-(4-hydroxyphenyl)-, mixt. with 4-(5-methyl-3-phenyl-4-isoxazolyl)benzenesulfonamide (CA INDEX NAME)

OTHER NAMES:

CN Valcox plus CN Valeron plus

MF C16 H14 N2 O3 S . C8 H9 N O2

CI MXS

SR CA

LC STN Files: CA, CAPLUS

CM 1

CRN 181695-72-7 CMF C16 H14 N2 O3 S

CM 2

CRN 103-90-2 CMF C8 H9 N O2

2 REFERENCES IN FILE CA (1907 TO DATE) 2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 7 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN

RN 935681-80-4 REGISTRY

ED Entered STN: 23 May 2007

CN B-Cyclodextrin, compd. with 4-(5-methyl-3-phenyl-4-isoxazolyl)benzenesulfonamide (1:?) (CA INDEX NAME)

FS STEREOSEARCH

MF C42 H70 O35 . x C16 H14 N2 O3 S

SR CA

LC STN Files: CA, CAPLUS

CM 1

CRN 181695-72-7

CMF C16 H14 N2 O3 S

CM 2

CRN 7585-39-9 CMF C42 H70 O35

Absolute stereochemistry.

PAGE 1-A

1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 8 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN

887258-63-1 REGISTRY RN

ED Entered STN: 08 Jun 2006

CN 2H-1, 2, 4-Benzothiadiazine-7-sulfonamide, 6-chloro-3, 4-dihydro-, 1,1-dioxide, mixt. with 4-(5-methyl-3-phenyl-4isoxazolvl) benzenesulfonamide (9CI) (CA INDEX NAME)

MF C16 H14 N2 O3 S . C7 H8 C1 N3 O4 S2

CI MXS

SR CA

LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL

CM 1

CRN 181695-72-7

CMF C16 H14 N2 O3 S

CM 2

CRN 58-93-5

CMF C7 H8 C1 N3 O4 S2

1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

ANSWER 9 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN RN

877140-46-0 REGISTRY ED

Entered STN: 17 Mar 2006

β-Cyclodextrin, compd. with 4-(5-methyl-3-phenyl-4isoxazolyl)benzenesulfonamide (1:1) (9CI) (CA INDEX NAME) FS STEREOSEARCH

MF C42 H70 O35 . C16 H14 N2 O3 S

SR CA LC STN Files: CA, CAPLUS

CM 1

CRN 181695-72-7

CMF C16 H14 N2 O3 S

CM 2

CRN 7585-39-9 CMF C42 H70 O35

Absolute stereochemistry.

PAGE 1-A



2 REFERENCES IN FILE CA (1907 TO DATE)
2 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L2 ANSWER 10 OF 162 REGISTRY COPYRIGHT 2008 ACS on STN

RN 862126-46-3 REGISTRY

ED Entered STN: 30 Aug 2005

CN Benzenesulfonamide, 4-[5-methyl-3-(2,4,6-trimethylphenyl)-4-isoxazolyl]-(CA INDEX NAME)

MF C19 H20 N2 O3 S

SR CA LC STN Files: CA, CAPLUS, CASREACT

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

1 REFERENCES IN FILE CA (1907 TO DATE)

1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> s 12 and valdecoxib?

4 VALDECOXIB?

3 L2 AND VALDECOXIB?

=> d 13 1-3

L3

- L3 ANSWER 1 OF 3 REGISTRY COPYRIGHT 2008 ACS on STN
- RN 676458-08-5 REGISTRY
- ED Entered STN: 22 Apr 2004
- CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, sodium salt (1:1) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, monosodium salt

```
(9CI)
OTHER NAMES:
CN Valdecoxib sodium
MF C16 H14 N2 O3 S . Na
SR CA
LC STN Files: CA, CAPLUS
CRN (181695-72-7)
```

Na

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L3 ANSWER 2 OF 3 REGISTRY COPYRIGHT 2008 ACS on STN RN 676458-07-4 REGISTRY

RN 676458-07-4 REGISTRY ED Entered STN: 22 Apr 2004

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, potassium salt (1:1) (CA INDEX NAME)

OTHER CA INDEX NAMES: CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, monopotassium salt (9CI)

OTHER NAMES: CN Valdecoxib potassium

MF C16 H14 N2 O3 S . K

SR CA

LC STN Files: CA, CAPLUS

CRN (181695-72-7)

● K

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1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)
```

```
L3 ANSWER 3 OF 3 REGISTRY COPYRIGHT 2008 ACS on STN
```

RN 181695-72-7 REGISTRY

ED Entered STN: 10 Oct 1996

N Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

OTHER NAMES:
CN 4-(5-Methyl-3-phenylisoxazol-4-v1)benzenesulfonamide

CN Bextra

CN SC 65872

CN Valdecoxib

CN Valecoxib

CN Valus CN Valz

MF C16 H14 N2 O3 S

CI COM

SR CA LC STN Files: ADISINSIGHT, ANABSTR, BIOSIS, BIOTECHNO, CA, CAPLUS,

CASREACT, CBNB, CHEMCATS, CSCHEM, EMBASE, HSDB*, IMSDRUGNEWS, IMSPATENTS, IMSPRODUCT, IMSRESEARCH, IPA, MRCK*, PATDPASPC, PHAR, PROUSDDR, PS, RTECS*, SYNTHLINE, TOXCENTER, USAN, USPAT2, USPATFULL (*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

837 REFERENCES IN FILE CA (1907 TO DATE)

33 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

844 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> s valdecoxib

L4 4 VALDECOXIB

=> d 14 1-4

T. 4 ANSWER 1 OF 4 REGISTRY COPYRIGHT 2008 ACS on STN

RN 676458-08-5 REGISTRY

ED Entered STN: 22 Apr 2004

CN

Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, sodium salt (1:1) (CA INDEX NAME)

OTHER CA INDEX NAMES:

Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, monosodium salt CN (9CI)

OTHER NAMES:

CN Valdecoxib sodium

MF C16 H14 N2 O3 S . Na

SR

LĊ STN Files: CA, CAPLUS

CRN (181695-72-7)

Na

1 REFERENCES IN FILE CA (1907 TO DATE) 1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

ANSWER 2 OF 4 REGISTRY COPYRIGHT 2008 ACS on STN

676458-07-4 REGISTRY RN

Entered STN: 22 Apr 2004

Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, potassium salt (1:1) (CA INDEX NAME)

OTHER CA INDEX NAMES:

Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, monopotassium salt (9CI)

```
OTHER NAMES:
CN Valdecoxib potassium
MF C16 H14 N2 O3 S . K
SR CA
LC STN Files: CA, CAPLUS
CRN (181695-72-7)
```

K

1 REFERENCES IN FILE CA (1907 TO DATE)
1 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L4 ANSWER 3 OF 4 REGISTRY COPYRIGHT 2008 ACS on STN

RN 181695-81-8 REGISTRY

ED Entered STN: 10 Oct 1996

CN Benzenesulfonamide, 4-[5-(hydroxymethyl)-3-phenyl-4-isoxazolyl]- (CA INDEX NAME)

OTHER NAMES:

CN 1-Hydroxyvaldecoxib

CN SC 66905

MF C16 H14 N2 O4 S

SR CA

LC STN Files: CA, CAPLUS, CASREACT, TOXCENTER, USPAT2, USPATFULL

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

- 54 REFERENCES IN FILE CA (1907 TO DATE)
- 3 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
- 54 REFERENCES IN FILE CAPLUS (1907 TO DATE)
- L4 ANSWER 4 OF 4 REGISTRY COPYRIGHT 2008 ACS on STN
- RN 181695-72-7 REGISTRY
- ED Entered STN: 10 Oct 1996
- CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)
- OTHER NAMES:
 CN 4-(5-Methyl-3-phenylisoxazol-4-yl)benzenesulfonamide
- CN Bextra
- CN SC 65872
- CN Valdecoxib
- CN Valecoxib
- CN Valus
- CN Valz
- MF C16 H14 N2 O3 S
- CI COM
- SR CA
- LC STN Files: ADISINSIGHT, ANABSTR, BIOSIS, BIOTECHNO, CA, CAPLUS, CASREACT, CBNB, CHEMCATS, CSCHEM, EMBASE, HSDB*, IMSDRUGNEWS, IMSPATENTS, IMSPRODUCT, IMSRESBEARCH, IPA, MRCK*, PATDPASPC, PHAR, PROUSDDR, PS, RIECS*, SYNTHLINE, TOXCENTER, USAN, USPATZ, USPATFULL (*File contains numerically searchable property data)

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

837 REFERENCES IN FILE CA (1907 TO DATE)
33 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
844 REFERENCES IN FILE CAPLUS (1907 TO DATE)

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION

225.17

224.96

FULL ESTIMATED COST

FILE 'CAPLUS' ENTERED AT 08:39:49 ON 23 JUN 2008
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FILE COVERS 1907 - 23 Jun 2008 VOL 148 ISS 26 FILE LAST UPDATED: 22 Jun 2008 (20080622/ED)

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http://www.cas.org/legal/infopolicy.html

=> s 14

DOCUMENT NUMBER:

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L5
          846 L4
=> s 15 and ?crystal?
        571556 ?CRYST
       2075241 ?CRYSTAL?
        571556 ?CRYST
       373142 CRYST
         1802 CRYSTS
       374411 CRYST
                 (CRYST OR CRYSTS)
        143102 ?CRYSTD
        94897 CRYSTD
        27466 ?CRYSTG
        20979 CRYSTG
        324049 ?CRYSTN
        251430 CRYSTN
          2463 CRYSTNS
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                 (CRYSTN OR CRYSTNS)
       2531614 ?CRYSTAL?
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                 OR CRYSTG OR ?CRYSTN OR CRYSTN)
           77 L5 AND ?CRYSTAL?
1.6
=> s 16 and polymorp?
        234268 POLYMORP?
            9 L6 AND POLYMORP?
=> d 17 1-9 ibib abs hitstr
L7 ANSWER 1 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                        2008:74254 CAPLUS <<LOGINID::20080623>>
```

TITLE: Compositions and methods comprising bicifadine for the treatment of chronic pain conditions

INVENTOR(S): Skolnick, Phil; Stern, Warren

PATENT ASSIGNEE(S): Dov Pharmaceutical, Inc., USA

SOURCE: PCT Int. Appl., 5pp.

148:175738

DOCUMENT TYPE: LANGUAGE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE			APPL	ICAT		DATE							
WO	WO 2008008474						A2 2008			117 WO 2007-US15964						20070711				
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,	CA,			
		CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,	FI,			
		GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,			
		KM,	KN,	KP,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	ME,			
		MG,	MK,	MN,	MW,	MX,	MY,	ΜZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,			
		PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	TJ,	TM,	TN,			
		TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	ZA,	ZM,	zw							
	RW:	ΑT,	ΒE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,			
		IS,	IT,	LT,	LU,	LV,	MC,	MT,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,			
		ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,			
		GH,	GM,	KΕ,	LS,	MW,	ΜZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,			
		BY,	KG,	ΚZ,	MD,	RU,	TJ,	TM												
US 20080014272							2008	0117	US 2007-775721						20070710					
PRIORITY APPLN. INFO.:									1	US 2	006-	8304	12P	1	P 2	0060	711			

B The present invention relates to methods, pharmaceutical compns, and kits for treating osteoarthritis-associated pain, inflammation and improving function in a patient comprising a first therapeutic agent which comprises bicifadine or a pharmaceutically acceptable salt, enantiomer, solvate, hydrate, polymorph or prodrug thereof and a second therapeutic agent which comprises a non-steroidal anti-inflammatory drug or derivative thereof. Thus, treatment with bicifadine or ibuprofen alone was no different than treatment with placebo in reducing osteoarthritis-associated pain as measured by the visual analog scale in humans. In contrast, treatment with the combination of bicifadine and ibuprofen resulted in a significant decrease in osteoarthritis-associated pain levels.

I 181695-72-7, Valdecoxib

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL

(Biological study); USES (Uses)
 (compns. and methods comprising bicifadine for treatment of chronic pain conditions)

RN 181695-72-7 CAPLUS

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

ACCESSION NUMBER: 2007:1483225 CAPLUS <<LOGINID::20080623>> DOCUMENT NUMBER: 148:449003

TITLE: Conformational aspects and interaction studies of heterocyclic drugs

AUTHOR(S): Ponnuswamy, M. N.; Gromiha, M. Michael; Sony, S. M.

Malathy; Saraboji, K.

CORPORATE SOURCE: Department of Crystallography and Biophysics, University of Madras, Chennai, 600 025, India
SOURCE: Topics in Heterocyclic Chemistry (2006), 3 (QSAR and Molecular Modeling Studies in Heterocyclic Drugs I),

81-147 CODEN: THCOA6; ISSN: 1861-9282

PUBLISHER: Springer GmbH

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review. Drug discoveries require the iterative synthesis-along with structural studies-of numerous individual analogs of biol. and medicinally active compds. Over half of all known compds. and a large number of pharmaceutical products are heterocyclic in nature. The pharmacol. activity of drugs depends mainly on interaction with their biol. targets, which have a complex three-dimensional structure, and mol. recognition is guided by the nature of the intermol, interactions. Furthermore, the drug's polymorphic nature also adversely affects its abilities. In order to address these factors, the stereochem. anal. of various piperidine and azepine derivs., weak π -interaction anal. of isoxazole, imidazole, indole, quinoline and triazole and polymorphic anal. of two com. drugs, valdecoxib and sildenafil citrate were carried out. Only the crystal structures were used for these analyses, of which the piperidine and azepine derivs., valdecoxib and sildenafil citrate were solved by our group. To understand the structure-activity relationship, the results of these studies were correlated with the crystal structure of their resp. drug mols. that are found in complex with the receptors. Stereochem, anal, showed that the ring conformation and orientation of the substituents correlate well with the active conformation of the drug. The π -systems prefer to form an offset stacking $\pi...\pi$ interaction geometry similar to the phenylalanine-phenylalanine interactions in proteins. polymorphic anal. one of the crystal conformations of valdecoxib proved to have better interaction with its receptor indicating higher activity.

IT 181695-72-7, Valdecoxib RL: PRP (Properties)

(conformational aspects and interaction studies of heterocyclic drugs) ${\tt RN} - 181695-72-7 - {\tt CAPLUS}$

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

L7 ANSWER 3 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:300882 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 147:350747

TITLE: Benzoquinolizine-2-carboxylic acid containing

compositions

INVENTOR(S): Saoji, Dilip Gopalkrishna; Nagori, Rajendra N.;

Saoji, Dilip Gopalkrishna; Nagori, Rajendra N.; Shukla, Milind Chintaman; Bhagwat, Sachin Subhash; Gupte, Shrikant Vinayak; Patel, Mahesh Vithalbhai;

Jha, Rasendrakumar; Kukreja, Anil; De Souza, Noel John Wockhardt Limited, India

PATENT ASSIGNEE(S): Wockhardt Limited, India SOURCE: Indian Pat. Appl., 34pp.

SOURCE: Indian Pat. Appl., 34p CODEN: INXXBQ

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
IN 2003MU01334	A	20060120	IN 2003-MU1334	20031231
PRIORITY APPLN. INFO.:			IN 2003-MU1334	20031231

AB This invention relates to topical compns. of an antibacterial benzoquinolizine-2-carboxylic acid, incorporated either as the single therapeutic ingredient in hitherto undescribed pharmaceutical compns., or as an ingredient in novel combination with at least one agent selected from a retinoid, an antifungal agent, another antibacterial compound and/or a steroid/non-steroid anti inflammatory agent, to processes for preparation of the compns., to use of the compns. in preparation of a medicament, and to a method of therapeutic or prophylactic use of such a composition for the treatment of dermal, ophthalmic, otic and nasal infections, with or without attendant inflammation.

IT 181695-72-7, Valdecoxib

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (benzoquinolizine carboxylic acid-containing topical compns.)

RN 181695-72-7 CAPLUS

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

L7 ANSWER 4 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:88439 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 146:169380

TITLE: Novel pharmaceutical modified release dosage forms comprising cyclooxygenase inhibitor

Jain, Rajesh; Jindal, Kour Chand; Singh, Sukhjeet;

Talwar, Munish PATENT ASSIGNEE(S):

Panacea Biotec Ltd., India PCT Int. Appl., 38pp. SOURCE: CODEN: PIXXD2

DOCUMENT TYPE: Pat.ent. LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

KIND DATE APPLICATION NO. PATENT NO. WO 2007010559 A2 20070125 WO 2006-IN258 A3 20070920 20060719 WO 2007010559 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,

AI, BE, BG, CH, CI, CZ, DE, DR, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, ML, PI, PIT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, MKE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA

IN 2005DE01899 A 20070824 IN 2005-DE1899 20070125 AU 2006-271150 A1 AU 2006271150 20060719 20070125 A1 20070125 CA 2006-2614850 A2 20080409 EP 2006-780539 CA 2614850 20060719 20060719 EP 1906933

R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA, HR, MK, RS

KR 2008032209 20080414 A PRIORITY APPLN. INFO.:

KR 2008-704111 IN 2005-DE1899 WO 2006-IN258

20080220 A 20050720 W 20060719

AB Pharmaceutical modified release dosage form comprising at least one cyclooxygenase inhibitor or its salts, esters, prodrugs, solvates, hydrates, or derivs. thereof as active agent, with a carrier for controlling the release of the cyclooxygenase enzyme inhibitor is provided. The dosage form preferably provides a release of not more than 60% of the cyclooxygenase enzyme inhibitor in 1 h and not less than 75% of the cyclooxygenase enzyme inhibitor after 12 h when tested in accordance with the dissoln, method using distilled water with 2.0% sodium lauryl sulfate as the dissoln. medium or in accordance with a dissoln. method employing pH 7.0 phosphate buffer with 2.0% sodium lauryl sulfate as the dissoln. medium or in accordance with a dissoln. method employing 0.001N HCl with 1.0% sodium lauryl sulfate as dissoln, medium. Further, the pharmaceutical composition of the present invention when tested in a group of healthy humans preferably achieves a mean peak plasma concentration (Cmax)

after

at least about 1 h of administration of the dosage form. The present invention also provides process of preparing such dosage form compns. and prophylactic and/or therapeutic methods of using such dosage forms. 181695-72-7, Valdecoxib

RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pharmaceutical modified release dosage forms comprising cyclooxygenase inhibitor)

181695-72-7 CAPLUS RN

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)



L7 ANSWER 5 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:1330315 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 144:57579

TITLE: Process for obtaining form A of valdecoxib suitable

for pharmaceutical formulations

INVENTOR(S): Thakashinamoorthy, Chandiran; Jesudoss, Mercy Gnanadeepam; Hariharasubramanian, Meera; Seetharaman,

Subramanian Sankara PATENT ASSIGNEE(S): India

SOURCE: PCT Int. Appl., 14 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	ENT :	NO.			KIN	D	DATE			APPL	ICAT:	DATE					
						_											
WO	2005	1204	99		A1		2005	1222	1	WO 2	004-	IN16:	2		2	0040	610
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,
		SI,	SK,	TR,	BF,	BJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,
		SN,	TD,	TG													
ORITY	APP	LN.	INFO	. :					1	WO 2	004-	IN16:	2		2	0040	610

PRIORITY APPLN. INFO.: WO 2004-IN162

AB The invention provides for a reactive crystallization procedure for obtaining polymorphic Form A of valdecoxib with desirable

particle size characteristics without milling, making it useful as an active ingredient in the preparation of pharmaceutical composition For example,

valdecoxib 250 g was added in to aqueous sodium hydroxide 5000 mL at 500 C and stirred. The content was heated to 60°C to get a clear solution The pH of the solution after dissoln. was 11.6. To the alkaline solution was added aqueous

hydrochloric acid 1280 mL through dip pipe. During the acidification the product ppts. from the solution of pH 1.5. The precipitated product was filtered

immediately and the product was then washed with water 6500 mL. The product obtained was dried at 70 to 75° C under reduced pressure till the water content in the product was less then 0.3 % weight/weight to vield

the product 235 g.

181695-72-7, Valdecoxib

RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES

(process for obtaining form A of valdecoxib suitable for pharmaceutical formulations)

RN 181695-72-7 CAPLUS

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

REFERENCE COUNT:

1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 6 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:823681 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 143:216704 TITLE:

Crystalline polymorphs of a

CXC-chemokine receptor ligand

INVENTOR(S): Hu, Mengwei; Yu, Younong; Dwyer, Michael; Taveras, Arthur G.; Kim-Meade, Agnes; Yin, Jianguo; Fu,

Xiaovong; Mcallister, Timothy; Zhang, Shuvi; Klopfer,

Schering Corporation, USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 65 pp. CODEN: PIXXD2

Patent

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE			
WO 2005075447	A1 20050818	WO 2005-US3414	20050128			
W: AE, AG, AL,	AM, AT, AU, AZ,	BA, BB, BG, BR, BW, BY,	BZ, CA, CH,			
CN, CO, CR,	CU, CZ, DE, DK,	DM, DZ, EC, EE, EG, ES,	FI, GB, GD,			
GE, GH, GM,	HR, HU, ID, IL,	IN, IS, JP, KE, KG, KP,	KR, KZ, LC,			
LK, LR, LS,	LT, LU, LV, MA,	MD, MG, MK, MN, MW, MX,	MZ, NA, NI,			
NO, NZ, OM,	PG, PH, PL, PT,	RO, RU, SC, SD, SE, SG,	SK, SL, SY,			
TJ, TM, TN,	TR, TT, TZ, UA,	UG, US, UZ, VC, VN, YU,	ZA, ZM, ZW			
RW: BW, GH, GM,	KE, LS, MW, MZ,	NA, SD, SL, SZ, TZ, UG,	ZM, ZW, AM,			
AZ, BY, KG,	KZ, MD, RU, TJ,	TM, AT, BE, BG, CH, CY,	CZ, DE, DK,			
EE, ES, FI,	FR, GB, GR, HU,	IE, IS, IT, LT, LU, MC,	NL, PL, PT,			

RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2005210504 20050818 AU 2005-210504 A1 20050128 CA 2554709 20050818 CA 2005-2554709 A1 20050128 US 20050192345 A1 20050901 US 2005-45772 20050128 EP 1723131 A1 20061122 EP 2005-712748 20050128 R: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LI, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, AL, BA, HR, LV, MK, YU CN 2005-80003507 CN 1914187 Α 20070214 20050128 BR 2005007329 Α 20070703 BR 2005-7329 20050128 JP 2007519751 Т 20070719 JP 2006-551613 20050128 MX 2006PA08599 Α 20060828 MX 2006-PA8599 20060728 IN 2006CN02800 20070608 IN 2006-CN2800 20060728 Α NO 2006003841 Α 20061027 NO 2006-3841 20060829 PRIORITY APPLN. INFO.: US 2004-540487P P 20040130 WO 2005-US3414 W 20050128

The present invention relates to 4 distinct crystalline polymorphs of a monohydrate of 2-hydroxy-N, N-dimethyl-3-[[2-[[1-(5methyl-2-furanyl)propyl]amino]-3,4-dioxo-1-cyclobuten-1vllamino|benzamide. These 4 polymorphic forms, herein referred to as Forms I, II, III and IV are active as a CXC-chemokine receptor ligands. The invention is further directed to formulations, methods of treatment, and processes of synthesis of these polymorphic forms.

- 181695-72-7, Valdecoxib
 - RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (crystalline polymorphs of CXC-chemokine receptor ligand)
- RN 181695-72-7 CAPLUS
- CN Benzenesulfonamide, 4-(5-methy1-3-pheny1-4-isoxazoly1)- (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

3 L7 ANSWER 7 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:565095 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 141:111581

TITLE: Benzoguinolizine-2-carboxvlic acid-containing

compositions

INVENTOR(S): Saoji, Dilip G.; Nagori, Rajendra N.; Shukla, Milind C.; Bhagwat, Sachin S.; Gupte, Shrikant V.; Patel, Mahesh V.; Jha, Rasendrakumar; Kukreja, Anil; De

Souza, Noel J.

PATENT ASSIGNEE(S):

India

SOURCE: PCT Int. Appl., 38 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

	PAT	TENT	NO.												DATE				
							_									-			
	WO	2004	0582	62		A1		2004	0715		WO 2	003-	IN42		20031231				
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
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			GE.	GH.	GM.	HR.	HU.	ID.	IL.	IN.	IS.	JP.	KE.	KG.	KP.	KR.	KZ.	LC.	
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	IN	2002																	
	CA	2512	190			A1		2004	0715		CA 2	003-	2512	190		2	0031	231	
	AU	2003	3023	05		A1		2004	0722		AU 2	003-	3023	05		2	0031	231	
	US	2004	0176	337		A1		2004	0909		US 2	003-	7499	33		2	0031	231	
	EP	1589	972			A1		2005	1102		EP 2	003-	8108	61		2	0031	231	
		R:	AT.	BE.	CH.	DE.	DK.	ES.	FR,	GB.	GR.	IT.	LT.	LU.	NL.	SE.	MC.	PT.	
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OTHER SOURCE(S): MARPAT 141:111581

AB The invention relates to topical compns. of an antibacterial

benzoquinolizine-2-carboxylic acid, incorporated either as the single therapeutic ingredient in hitherto undescribed pharmaceutical compns., or as an ingredient in novel combination with at least one agent selected from a retinoid, an antifungal agent, another antibacterial compound and/or a steroidal/nonsteroidal anti-inflammatory agent. Processes for preparation of the compns., the use of the compns. and a method of therapeutic or prophylactic use of such a composition for the treatment of dermal, ophthalmic, otic and nasal infections, with or without attendant inflammation are disclosed. Thus, a gel contained RS-(+)-9-fluoro-6,7-dihydro-8-(4-hydroxypiperidin-1-yl)-5-methyl-1-cxo-1H,5H-benzo[i,j]quinolizine-2-carboxylic acid 1.00, Carbopol 1.20, NaOH 0.112, diethanolamine 0.36, discodium dedetate 0.10, sodium sulfite 0.05, and water qs to 100%.

IT 181695-72-7, Valdecoxib

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (benzoquinolizinecarboxylic acid-containing topical compns.)

RN 181695-72-7 CAPLUS

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

L7 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2008 ACS on SIN

ACCESSION NUMBER: 2004:269999 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 140:309372

TITLE: Pharmaceutical compositions with improved dissolution INVENTOR(S): Remenar, Julius; Peterson, Matthew, Almarsson, Om; Guzman, Hector; Chen, Hongming; Tawa, Mark; Oliveira,

Mark
PATENT ASSIGNEE(S): Transform Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 185 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 18
PATENT INFORMATION:

PAT	ENT :	NO.			KIND DATE					ICAT			DATE						
					A2 20040401 A3 20040805								20030916						
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		FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PT,	RO,	SE,	SI,	SK,	TR,		
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AU	2003	2672	31					0408		AU 2	003-	2672	31		2	0030	916		
CA	2511	881			A1		2004	0722		CA 2	003-	2511		2	0031	224			
WO	2004	0614	33		A1		2004	0722		WO 2	003-	US41		2	0031	224			
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ΑU	2003	3303591 A1 20040729						0729		AU 2	003-	3035	91		20031224				
EP	1579	79198 A1 20050928						0928		EP 2	003-	8085	67		20031224				
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IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
    JP 2006517527
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    WO 2004060347
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                                         WO 2003-US41642
                                                                20031229
                        A2
    WO 2004060347
                        A3
                              20041104
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
            CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE,
            GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
            LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
            OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
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    AU 2003300452
                       A1 20040729 AU 2003-300452
                                                              20031229
    US 20060052432
                        A1
                              20060309
                                         US 2005-528244
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    US 20060134198
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    US 20060140985
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                                         US 2005-541703
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PRIORITY APPLN. INFO.:
                                          US 2002-412459P
                                                           P 20020920
                                          US 2002-426275P
                                                            P 20021114
                                          US 2002-427086P
                                                            P 20021115
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                                                            P 20021126
                                          US 2002-437516P
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                                                             P 20030318
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                                                               20020903
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                                                            A 20030530
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                                                            A 20030620
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                                                            A 20030620
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                                                            P 20030711
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                                                            A 20030808
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                                                            A 20030911
                                          WO 2003-US28982
                                                            W 20030916
                                          WO 2003-US41273
                                                            W 20031224
                                          WO 2003-US41642
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AB The invention relates to methods of screening mixts. containing a pharmaceutical compound and an excipient to identify properties of the pharmaceutical compound/excipient combination that retard solid-state nucleation. The invention further relates to increasing the solubility, dissoln. and bioavailability of a drug with low solubility in gastric fluids conditions by combining the drug with a precipitation retardant and an optional enhancer. For example, celecoxib sodium salt was prepared from 126.3 mg of celecoxib in isopropanol and sodium enthoxide (21% ethanol solution). Water was added to a 1:4 mixture of celecoxib sodium salt and polyvinylpyrrolidone to obtain a clear solution The solution was stable for at least 15 min, after which time, crystals of neutral celecoxib began to form.
Crystalline neutral celecoxib do to dissolve when added to aqueous

Crystalline neutral celecoxib did not dissolve when added to aqueous polyvinylpyrrolldone or when water was added to a dry blend of neutral crystalline celecoxib and polyvinylpyrrolidone.

IT 181695-72-7, Valdecoxib 676458-07-4, Valdecoxib potassium 676458-08-5, Valdecoxib sodium

RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(compns. with improved dissoln. and bioavailability of drugs with low solubility)

- RN 181695-72-7 CAPLUS
- CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)

- RN 676458-07-4 CAPLUS
- CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)-, potassium salt (1:1) (CA INDEX NAME)

● K

- RN 676458-08-5 CAPLUS

● Na

L7 ANSWER 9 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER:

2002:107158 CAPLUS <<LOGINID::20080623>>

DOCUMENT NUMBER: 136:161365

TITLE: Aldosterone antagonist-cyclooxygenase-2 inhibitor

combination therapy to prevent or treat inflammation-related cardiovascular disorders

INVENTOR(S): Rocha, Ricardo; Zack, Marc D.; McMahon, Ellen G. PATENT ASSIGNEE(S):

Pharmacia Corporation, USA SOURCE: PCT Int. Appl., 273 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

					KIND DATE							DATE					
					A2 20020207 A3 20021128				WO 2	001-		20010726					
	W:	CO, GM, LS, RO,	CR, HR, LT, RU,	CU, HU, LU, SD,	CZ, ID, LV,	DE, IL, MA, SG,	DK, IN, MD,	AZ, DM, IS, MG, SK,	DZ, JP, MK,	EC, KE, MN,	EE, KG, MW,	ES, KP, MX,	FI, KR, MZ,	GB, KZ, NO,	GD, LC, NZ,	GE, LK, PL,	GH, LR, PT,
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CA	2415	078			A1		2002	0207		CA 2	001-	2415	078		2	0010	726
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EP	1303	308			B1		2006	0906									
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US	2003	0125	312		A1		2003	0703		US 2	001-	9157	84		2	0010	726
US	6716	829			B2		2004	0406									
JP	2004	5050	60		T		2004	0219		JP 2	002-	5153	11		2	0010	726
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	2469										001-						
	2002										002-						
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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR BR 2001017195 BR 2001-17195 Α 20050209 20011213 CN 1582154 20050216 CN 2001-823946 А 20011213 JP 2005516015 T 20050602 JP 2003-552305 20011213 US 20030191100 20031009 US 2002-243876 20020913 A1 MX 2004PA05803 A MX 2004-PA5803 20041101 20040614 US 20070191324 A1 20070816 US 2006-613879 20061220 PRIORITY APPLN. INFO.: US 2000-221364P P 20000727 US 2001-261497P P 20010112 US 1999-164390P P 19991109 US 2000-211064P P 20000613 US 2000-211250P P 20000613 US 2000-211253P P 20000613 US 2000-211264P P 20000613 US 2000-211311P P 20000613 US 2000-211340P P 20000613 US 2000-211451P P 20000613 US 2000-211459P P 20000613 US 2000-221358P P 20000727 US 2000-233056P P 20000914 US 2000-709253 A2 20001108 US 2000-712543 A1 20001114 US 2000-713348 B2 20001114 US 2001-261352P P 20010112 WO 2001-US23601 W 20010726 WO 2001-US48419 W 20011213

OTHER SOURCE(S): MARPAT 136:161365

AB Combinations of aldosterone blockers and Cyclooxygenase-2 inhibitors useful in the treatment of inflammation-related cardiovascular disorders are disclosed.

IT 181695-72-7

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(aldosterone antagonist-cyclooxygenase-2 inhibitor combination therapy to prevent or treat inflammation-related cardiovascular disorders)

US 2003-682527

A1 20031009

RN 181695-72-7 CAPLUS

CN Benzenesulfonamide, 4-(5-methyl-3-phenyl-4-isoxazolyl)- (CA INDEX NAME)